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A 64-Nuclear Cubic Cage Incorporating Propeller-like Fe^{III}₈ Apices and HCOO⁻ Edges

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High nuclearity clusters are currently of great interest, by virtue of their structural beauty and their important role in connecting micro to macro and quantum to classic regimes.¹⁻¹⁰ While many such systems have been explored, for example, giant polyoxometalates,1 giant metal-chalcogenide clusters,2 inorganic fullerene-like molecules,³ lanthanide hydroxo compounds,⁴ and oxo-clusters containing both d- and f-block metals,⁵ those containing magnetic transition metals attract special and continuous attention regarding their rich magnetism such as high ground-state spin values,⁶ single molecular magnet (SMM)⁷ and coupled SMM behavior by weak linkages,⁸ and their promising applications for data storage,⁷ quantum computing,⁹ and spintronics.¹⁰ Various ligands, especially multipodal ones,¹¹ are widely employed in the syntheses of such high nuclearity species. However, many simple and popular ligands, like acetate, can also afford compounds of high nuclearity.¹² With these serendipities in mind, we are currently exploring the mixedligand system of triethanolamine (N(CH₂CH₂OH)₃, teaH₃), a tripodal ligand having been recently used for synthesizing polynuclear species,^{11,13} and formate HCOO⁻, the smallest and simplest carboxylate of small stereohindrance, favoring both formation of metal-oxo clusters and further linkage of them.¹⁴ The outcome is, quite surprising and interesting, a large cubic cage consisting of 64 Fe^{3+} ions of {[Fe₈O₃(*tea*)(*tea*H)₃(HCOO)₆]₈(HCOO)₁₂}- $(ClO_4)_{12} \bullet 3CH_3OH \bullet 36H_2O (Fe_{64}) (tea = N(CH_2CH_2O)_3^{3-} and teaH$ = $N(CH_2CH_2O)_2$ -(CH_2CH_2OH)²⁻). Fe³⁺-oxo-based clusters have been continuously studied.^{15–18} They can display ground state of spin values from 0 up to 25 and pleasing physical properties such as the Néel vector tunneling of Fe₈ wheel¹⁷ and the large magnetocaloric effect in an Fe₁₄ cluster.¹⁸

Fe₆₄ was obtained by a very slow crystallization procedure or reaction lasting 2 months (Supporting Information, SI). The structure analysis (SI) revealed a cubic cage (Figure 1) containing 64 Fe³⁺ ions, 24 O²⁻, 8 tea³⁻, 24 teaH²⁻, and 60 HCOO⁻, thus a total of 180 components. The cubic cage consists of eight corners of octanuclear [Fe^{III}₈O₃(*tea*)(*tea*H)₃(HCOO)₆(HCOO)_{3/2}] unit (Fe₈, Figure 1a,b) with a propeller-like Fe₈O₃ core, linked further by 12 anti-anti formato ions as the edges of the cube (Figure 1c). The Fe₈ corner possesses similar structure to [Fe₈O₃(O₂CPh)₉-(tea)(teaH)₃] • MeCN and related octanuclear compounds,¹⁶ and the Fe₈ moiety has been found in some high nuclearity clusters.^{15b} Its propeller-like core has two central axial Fe³⁺ ions bridged by three μ_4 -O²⁻, and each μ_4 -O²⁻ further bridges two peripheral Fe³⁺ ions in pair, one upper and one lower (Figure 1b), being one of the three blades of the propeller. The Fe₈O₃ core thus consists of three flattened [Fe₄(μ_4 -O)] tetrahedra sharing the central Fe–Fe axis of the core. The Fe^{3+} ions are further linked by ligands, tea^{3-} , $teaH^{2-}$, and HCOO⁻. The upper axial Fe^{3+} ion is capped by one *tea*³⁻ via the N atom and the three alkoxide O atoms, thus this Fe³⁺ ion is 7-fold coordinated in a face-capped octahedron. Each alkoxide O



Figure 1. The structure of Fe₆₄ cubic cage. (a and b) Top and side view of the Fe₈ corner. Atomic scheme: Fe, large green (peripheral) and yellow (central) spheres; O, small red and purple (the three μ_4 -O²⁻) spheres; N, blue spheres; C, gray spheres. (c) The cubic cage with the corner Fe₈O₃ cores and HCOO⁻ edges highlighted (O red and C black) while other parts in thin sticks. H atoms, ClO₄⁻ anions, and lattice solvents are not included.

atom of the tea^{3-} connects one upper peripheral Fe³⁺ ion. However, each lower peripheral Fe^{3+} ion is chelated by one *tea*H²⁻ through its N atom and two alkoxide arm O atoms, of which one further links the lower central Fe³⁺ ion, and the other to one upper peripheral Fe³⁺ ion, leaving one alcohol arm free of coordination. Therefore, the lower axial Fe³⁺ ion is octahedrally coordinated by O atoms. In each blade, the two peripheral Fe³⁺ ions are bridged by two syn-syn HCOO⁻ ligands, and the upper peripheral Fe³⁺ ion is finally coordinated by one HCOO⁻ ligand, which is one edge connecting to other Fe₈ corner of the cube (Figure 1b,c). Each Fe₈ corner thus has three HCOO⁻ ligands to link the other three Fe₈ corners, and its central axis points to the cube center, with the tea^{3-} cap inside the cube. It is noted that the coordinate octahedra of Fe³⁺ ions are face-sharing for the two central axial Fe³⁺ ions (Fe···Fe 2.994 Å), edge-sharing between the central and peripheral Fe³⁺ ions (Fe···Fe 2.988-3.132 Å), apex-sharing between the central and peripheral Fe³⁺ ions (Fe···Fe 3.638-3.902 Å), and apex-sharing between the peripheral Fe^{3+} ions (Fe···Fe 3.270-3.539) Å). It could be estimated that the magnetic interaction within the Fe8 corner should be strong or major. The Fe···Fe distances spanned by the anti-anti HCOO⁻ edge are 5.806-5.943 Å, and the cube has edges of 13.2 Å counting the separation between the outer axial Fe atoms of the corners. It is clear that the small stereohindrance of HCOO⁻ ligand¹⁴ allows the further linkage of the bulk Fe₈ units. The successful synthesis of the Fe₆₄ cage reveals that the HCOO⁻ ligand is indeed benefitial for synthesizing species of high nuclearity. In the lattice, the cationic Fe₆₄ cubic cages have their 3-fold axes parallel to the c axis, and form the hexagonal



Figure 2. χT versus T plots under 1 kOe field, and M versus H plots (inset) for Fe_{64} and Fe_{8a} . Data are for one Fe^{3+} .

closest packing layer (Figure S1 in SI) along the ab plane, then these layers stack along the *c* direction in ABCABC style, with interstices occupied by ClO₄⁻ anions, lattice water and methanol, mostly disordered. The shortest intercage Fe····Fe separation is 8.61 Å.

In a different preparation using Fe³⁺ source, we obtained the compound of [Fe₈O₃(tea)(teaH)₃(HCOO)₉]{[(CH₃CH₂)₃NH+]- $(ClO_4^{-})_{3}(H_2O)_{0.5}$ (**Fe**₈**a**), in quite a high yield (SI). The compound consists of discrete neutral octanuclear [Fe₈O₃(tea)(teaH)₃(HCOO)₉] molecules, with similar molecular structure (Figure S2 in SI) to the Fe₈ corner of the Fe₆₄ cage. The shortest intercluster Fe····Fe separation is 7.05 Å. However, several attempts to prepare Fe_{64} from Fe³⁺ source were unsuccessful (SI).

The magnetic study of Fe₆₄ was performed on carefully selected small crystals of Fe₆₄ but still containing a very small amount of impurity of the ferrimagnetic [Fe3(HCOO)6](solvent)19 and weak ferromagnetic Fe(HCOO)₂•2H₂O²⁰ probably (SI). As depicted in Figure 2, at 300 K, the χT value is 1.39 cm³ mol⁻¹ K per Fe³⁺. significantly lower than the value of 4.375 cm³ mol⁻¹ K expected for the noninteracting high-spin Fe^{3+} ion (S = 5/2 and g = 2.00). It decreases almost linearly with lowering temperature to 0.016 cm³ mol^{-1} K at 2.0 K. A broad maximum of the corresponding χ is observed at 170 K (Figure S3 in SI). The overall behavior is clearly indicative of strong antiferromagnetic interactions between the Fe³⁺ centers. The rise in χ values and a kink in χT around 16 K is due to the impurity of ferrimagnetic [Fe3(HCOO)6](solvent) with critical temperature of 16.1 K,¹⁹ and further rise in χ values below 5 K might be due to the impurity of weak ferromagnetic Fe(HCOO)₂·2H₂O with critical temperature of 3.7 K.²⁰ The impurity is responsible for the rapid rise of the isothermal magnetization in the low field region at 2 K (Figure 2 inset). However, the very low value of 0.08 N β per Fe³⁺ even at 50 kOe, far below the saturation value expected (5 N β per Fe³⁺ assuming g = 2.00), confirms the significant antiferromagnetic interactions within the material. The isothermal magnetization at 30 K, above the critical temperatures of the two assumed impurities, is linear and reaches only 0.06 N β per Fe³⁺ at 50 kOe. The magnetic behavior of the Fe₆₄ cage is similar to the related discrete octanuclear species¹⁶ possessing the same propeller-like Fe₈O₃ core. It is clear that in the Fe₆₄ cage the antiferromagnetic interaction within the Fe₈ corner is strong and dominant, and the magnetic interaction between Fe₈ corners through the anti-anti HCOO⁻ edge is minor but it could have an effect on the magnetism. The higher peak temperature of 170 K of Fe₆₄ versus 80 K of Fe₈a in temperature-dependent susceptibility (Figure S3 in SI) revealed that the anti-anti HCOO⁻ bridges in the Fe₆₄ cage provide further significant interaction between the Fe8 corners, compared to Fe8a which contains discrete neutral octanuclear species only. Both Fe₆₄ and $Fe_{8}a$ have S = 0 ground state as observed for many evennumbered Fe³⁺-oxo-based clusters.^{15,16}

In conclusion, a high nuclearity cubic cage involving 64 Fe^{3+} ions has been successfully synthesized, and it displays strong antiferromagnetism. This demonstrates that the combination of the small stereohindrance HCOO⁻ and polypodal ligands can provide a new route to obtain high nuclearity magnetic clusters and/or further linkage of them, thus giving the opportunity to explore their beautiful structures and interesting magnetism.

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Supporting Information Available: An X-ray crystallographic file in CIF format for Fe₆₄ and Fe₈a, a PDF file containing further information and Figures S1-S3. This material is available free of charge via the Internet at http://pubs.acs.org.

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